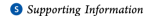


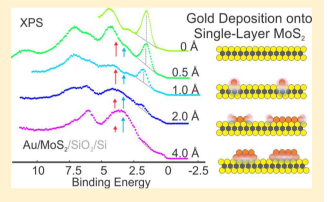
# Gold Dispersion and Activation on the Basal Plane of Single-Layer

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ABSTRACT: Gold islands are typically associated with high binding affinity to adsorbates and catalytic activity. Here we present the growth of dispersed nanoscale gold islands on single layer MoS<sub>2</sub>, prepared on an inert SiO<sub>2</sub>/Si support by chemical vapor deposition. This study offers a combination of growth process development, optical characterization, photoelectron spectroscopy at submicron spatial resolution, and advanced density functional theory modeling for detailed insight into the electronic interaction between gold and single-layer MoS2. In particular, we find the gold density of states in Au/MoS<sub>2</sub>/SiO<sub>2</sub>/Si to be far less well-defined than Au islands on other 2-dimensional materials such as graphene, for which we also provide data. We attribute this effect to the presence of heterogeneous Au adatom/MoS<sub>2</sub>-support interactions within the nanometer-scale gold cluster. Theory



predicts that CO will exhibit adsorption energies in excess of 1 eV at the Au cluster edges, where the local density of states is dominated by Au 5d<sub>2</sub>2 symmetry.

#### INTRODUCTION

Transition metal dichalcogenides (TMDs) such as MoS<sub>2</sub> have many and diverse technological applications ranging from chemical catalysis to emerging microelectronic integration to lubrication of surfaces. In particular, MoS<sub>2</sub> is the key industrial catalyst for hydrodesulfurization of crude oil; other catalytic applications of MoS2 include the formation of alcohols from syngas,<sup>2</sup> as a reverse water-gas shift catalyst,<sup>3,4</sup> and for facilitation of the hydrogen evolution reaction in electrocatalysis.<sup>4,5</sup> While oxide-based catalytic compounds are frequently enhanced by metal nanoparticles, MoS<sub>2</sub> single-layer islands support a metallic state at their edge, to which catalytic properties have been attributed.<sup>6</sup> At the same time, gold nanoparticles have demonstrated catalytic properties<sup>7-9</sup> different from bulk gold that are substrate- and particle-size dependent. On the archetypical 2-dimensional van der Waals material, graphene, gold aggregates even at very low dosages to

form a bulk-like electronic structure that is catalytically inactive. 10,11 Substantial substrate interactions are required to prevent surface aggregation of this most noble of elements; 12 reducible oxide surfaces such as titania or ceria are typically used for gold dispersion. 13,14

Here we investigate the binding of gold to a single-layer MoS<sub>2</sub> film on an oxide support and explore the shape/ electronic properties of the resulting gold overlay. The goal is to understand the gold-MoS2 substrate interactions, followed by the electronic structure of the resulting nanocluster, and their catalytic activity as a function of size. 15-17 Gold nanoparticles have been investigated on MoS<sub>2</sub>, <sup>18</sup> for instance for electrode applications 19 and as a target for optical control of

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particle properties. <sup>20</sup> We find that the MoS<sub>2</sub> van der Waals basal plane differs substantially from graphene: gold is dispersed into clusters small enough to exhibit an electronic structure typical of catalytically active gold including a shift/broadening of the d-band position.

Our findings confirm the computational predictions of Rawal et al. <sup>21</sup> that suggest Au nanoparticles to bind strongly to single-layer MoS<sub>2</sub>; Rawal et al. <sup>21</sup> underscore the potential of the Au/MoS<sub>2</sub> materials system for catalytic application. Furthermore, unlike several prior studies that define the impact of the substrate on the catalytic activity as indirect, i.e., caused by preventing aggregation but not by specific chemical gold-substrate interactions, <sup>22,23</sup> our computational studies trace the potential catalytic activity of the Au/MoS<sub>2</sub> system to the emergent electronic structure of the composite. We note that recent studies of Au on h-BN<sup>24–26</sup> also suggest substantial Au island to substrate interactions.

There is a resemblance between  $MoS_2$  particles found in actual catalysts<sup>27</sup> and the single-layer islands we can prepare efficiently on an otherwise inert substrate. The availability of the latter on a flat substrate renders them better accessible to spatially and angular resolved photoelectron spectroscopy while not compromising the relevance of our results. We find that the  $MoS_2$  layer acts as a facile gold-dispersion interlayer. In this regard, our approach offers a direct counterpoint to prior work by Besenbacher and Lauritsen's group, in which  $MoS_2$  was prepared on bulk gold and its properties investigated.<sup>6</sup>

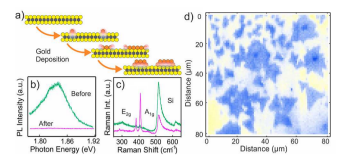
Single-layer MoS<sub>2</sub> can be grown by CVD on a broad range of substrates to coat them in a near-conformal fashion offering scalable access to the material system investigated here. MoS<sub>2</sub> single- (or few-)layer coating leading to Au-dispersion and -activation on a comparatively inert substrate like SiO<sub>2</sub>, that otherwise does not generally support gold dispersion, has potential appeal for extending the use of gold nanoparticle catalysts to a broader range of support materials with unique thermal, mechanical, and porosity properties.

Single- and few-layer TMD films have recently attracted intense interest for electronic applications as semiconducting 2D counterparts of graphene. A large number of photoelectron spectroscopy studies have addressed in this context the band structure of  $MoS_2$  both in the bulk and thinned (generally by mechanical exfoliation) to the single-layer limit. These provide us with a reference ascertaining that the CVD  $MoS_2$  material employed in this study exhibits the anticipated band structure near  $\Gamma$  (see the Supporting Information, SI). Analysis of the Fermi level position of our material ascertains the low sulfur vacancy defect density. We note that in prior studies we addressed the activation of  $MoS_2$  by sulfur depletion  $MoS_3$ —and do not see any such effects in this study.

This study combines chemical vapor deposition (CVD) growth of single-layer single-crystalline  ${\rm MoS_2}^{34}$  on ultrathin  ${\rm SiO_2}$  on Si substrates, high-resolution synchrotron-based spectro-imaging and ultraviolet photoelectron spectroscopy, and density functional theory (DFT) modeling to understand the binding of gold to the basal plane of single-layer  ${\rm MoS_2}$  as a function of gold coverage (Figure 1a).

# METHODS

Chemical Vapor Deposition Growth of MoS<sub>2</sub> Islands. This work employs chemical vapor deposition (CVD) molybdenum disulfide (MoS<sub>2</sub>) single-layer single-crystalline islands grown onto a  $\sim$ 1 nm SiO<sub>2</sub> film supported by a silicon wafer. CVD growth follows ref 34 and proceeded in a quartz



**Figure 1.** a) Schematic representation of the experiment and results: as the Au coverage on single-layer  $MoS_2$  islands increases from isolated atoms to single- and bi-layer islands, strong Au-MoS $_2$  interaction at isolated atoms, island edges, and the bottom layer modify the Au 5d density of states. The layer thickness/island density is for illustrative purposes and does not reflect the experimental values quantiatively; b) photoluminescence (PL) spectroscopy shows a bright signal at the optical bandgap of  $MoS_2$  prior to gold deposition and complete quenching following deposition of 4 Å of gold; c) Raman spectroscopy reveals surface enhanced  $MoS_2$   $E_{2g}$  and  $A_{1g}$  signal once the gold is deposited (at unchanged position) as well as attenuation of the Si substrate peak; and d) map of the lateral variation of the photoelectron yield reveals the size and location of the single-layer single-crystalline  $MoS_2$  islands used in this study.

tube furnace utilizing molybdenum trioxide (MoO<sub>3</sub>, Aldrich) and elemental sulfur (Aldrich) as precursors. Inert nitrogen gas was used to transfer sulfur vapor from an alumina boat held slightly above the sulfur's melting temperature to the hot (650 °C) reaction zone at the tube furnace center, where the target substrate was supported from a second alumina boat that also contained a small quantity (50 mg) of MoO<sub>3</sub>. Single-layer MoS<sub>2</sub> island deposition and ripening occurred over 10 min at the maximum furnace temperature followed by a slow cool-down.

Optical Characterization of Single-Layer MoS<sub>2</sub> Material. Optical characterization utilizes a Horiba LabRAM system and a laser at a wavelength of 532 nm. Because of the absence of sizable oxide layer, the single-layer MoS<sub>2</sub> islands are barely revealed in optical imaging. Photoluminescence and Raman spectroscopy utilizes an excitation power of 0.1 mW focused on a diffraction limited spot. The same sample area was investigated prior to and after gold deposition.

Photoelectron Spectroscopy and Gold Deposition. The ARPES experiments were performed at the 3.2L spectromicroscopy undulator beamline of the Elettra light source at a temperature of 99 K using a photon energy of 27 eV. The incident radiation was linearly polarized (along the horizontal direction) and focused on a 0.6  $\mu$ m diameter spot by means of a Schwarzschild objective.<sup>2</sup> An incident angle of 45° with respect to the sample surface was used to optimize surface sensitivity. The ARPES data were acquired using a hemispherical electron energy analyzer with a combined energy resolution of 50 meV and angular resolution of <1°. The sample was mounted onto a scanning stage, which enabled positioning and raster imaging with respect to the fixed photon beam. Prior to photoelectron spectroscopy, the sample was annealed to 250 °C to remove any spurious water adsorption. Gold deposition took place in situ inside the same ultrahigh vacuum system at a rate of 0.05 Å/s using a thermal evaporator and a quartz crystal microbalance for calibration.

**Computational Approach.** Our DFT simulations employ the projector-augmented wave (PAW)<sup>38,39</sup> and plane wave basis set methods. We used the Perdew–Burke–Ernzerhof

functional (PBE)40 to describe the electronic exchangecorrelation together with the DFT-D3 correction<sup>41</sup> for accounting wan der Waals interactions. The energy cutoff for plane-wave expansion was set at 500 eV, which is sufficient for energy and force convergences. The simulation supercell consists of an 8 × 8 single layer MoS2 constructed using the optimized lattice parameter (a = 3.16 Å), an Au cluster adsorbed on one side of the MoS, layer and a vacuum of about 15 Å to separate the resultant structure with its normal periodical image. Because of the large supercell, we sample the Brillouin Zone with one point locating at the zone center for structural relaxation (with residual force of less than 0.01 eV/Å) and a  $3 \times 3 \times 1$  k-point mesh centered at the zone center for density of states (DOS) calculations. For visualization purposes, we use a Gaussian filter with  $\sigma$  = 0.1 eV to obtained DOS. Since the structures of the Au clusters are not resolved in experiment, and unknown, it is impossible to have one-to-one correspondence between experiment and simulation. Thus, for demonstration purpose, we choose Au clusters of 1, 7, 19, and 31 atoms, representing the smallest, two hexagonal-one-layer, and one hexagonal-bilayer clusters, respectively. For each cluster, we survey several adsorption configurations on MoS2 and we find that the lowest energy one is the one in which the center Au atom is on top of an S atom of MoS2. The DOS analyzed and presented in this work are those of the lowest energy configurations.

# RESULTS

The material used in this study shows the expected bright photoluminescence (PL) and Raman features of single-layer MoS<sub>2</sub> (Figure 1b,c). <sup>28,29,34</sup> The photoluminescence is slightly red-shifted as typically found in the absence of an SiO<sub>2</sub> buffer layer. The MoS<sub>2</sub> Raman features are less pronounced than on, for instance, SiO<sub>2</sub> due to substrate screening.<sup>42</sup> We observe the  $E_{2g}$  and  $A_{1g}$  modes are found at is 383 and 408 cm<sup>-1</sup>, respectively, and attribute the comparatively wide Raman mode separation to the same substrate interactions that cause strong substrate adhesion to bare Si. Notably, the small mode separation of single layer material is due to the absence of bonding to layers above and below. The case of an interacting substrate, like the silicon used in this study, differs from inert oxide typically used in studies of single-layer MoS2. It is not astonishing that the out-of-plane vibrational  $A_{1g}$  mode is hardened in the presence of this interacting substrate, while the in-plane  $E_{2g}$  mode remains at the value expected for single-layer material.

We find significant enhancement of the  $MoS_2$  Raman signal after deposition of 4 Å of gold indicative of the presence of very small gold islands that support plasmon-enhanced Raman measurements<sup>43</sup> or surface enhance Raman (SERS). Complete quenching of the PL signal after gold deposition at low average coverage indicates high dispersion of the gold (either as a continuous film or into small clusters) and absence of a combination of large-scale aggregation and empty areas (which would retain their PL signal). We note that a layered (Frank van der Merve) growth of Au on  $MoS_2$  can be ruled out given the high surface energy of gold and the comparatively low growth temperatures; thus, the efficient quenching of the  $MoS_2$  PL signal observed in this study directly indicates the formation of highly dispersed small-size gold aggregates.

Mapping of the photoelectron yield across the substrate reveals that the substrate is covered with an array of single-crystalline triangular  $MoS_2$  islands of  $\sim 10~\mu m$  in size

interspersed between islands that originate from more than one nucleation spot and, consequently, contain a limited number of internal rotational domains (Figure 1d). Optical imaging of the islands on the substrate offers very low contrast because of the absence of a sufficient oxide layer underneath the single-layer  $MoS_2$  islands. <sup>44</sup>

Figure 2a shows a series of photoelectron spectra taken at the same spot as the Raman and PL measurements of Figure 1b,c

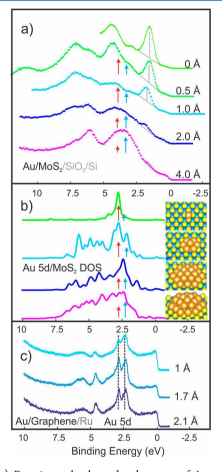


Figure 2. a) Experimental valence band spectra of Au on the same spot of an island of single-layer MoS<sub>2</sub>/SiO<sub>2</sub>/Si. With increasing gold coverage (~2.5 Å correspond to a monolayer of gold), the Au-derived signal dominates the spectrum and shifts from higher (red arrow) to lower (blue arrow) binding energy while the MoS<sub>2</sub> VB intensity (dashed line) is reduced and broadened by Au-MoS2 interactions; b) computational Au 5d density of state of the four clusters indicated. As the cluster size increases, lower binding energy gold DOS (blue arrow) augments and ultimately overshadowed the higher binding energy DOS (red arrow) found for a single Au atom. As these calculations do not include the SiO<sub>2</sub>/Si substrate and any ensuing surface band bending, the Fermi level position (x-axis) was adjusted for optimal match with the experimental data; and c) experimental reference spectra of Au on graphene show a clear Au 5d doublet and pronounced Fermi edge indicative of large metallic island formation and low substrate interaction, in great contrast to Au/MoS<sub>2</sub>.

on a single-layer  $MoS_2$  islands (shown in Figure 1d) as the gold coverage was successively increased from 0 to 0.5, 1, 2, and 4 Å (approximately 0, 0.2, 0.4, 0.8, and 1.6 monolayers, respectively). The final average coverage falls slightly short of a bilayer of gold. The angle-integrated spectra are based on angular-resolved measurements that at no and low gold coverage exhibit the known  $MoS_2$  dispersion near the  $\Gamma$  point

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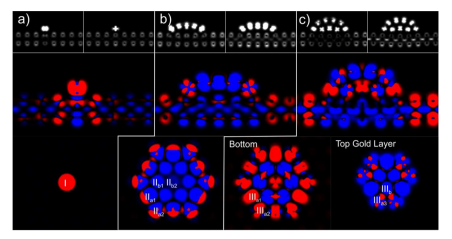


Figure 3. Comparison of the distribution of the density of state in the energy intervals marked by the red and blue arrows of Figure 2 (top row, left and right respective) and represented as difference for a) a single gold atom, b) a 19 atom cluster, and c) a bilayer 31 atom cluster of gold on  $MoS_2$ . The  $MoS_2$ -affected gold DOS is consistently found at the perimeter of the cluster suggesting a propensity for activity of this particular region. Test sites for CO adsorption are labeled by roman numerals followed by a and b for a site at which the higher or lower binding energy state dominates, respectively.

(see the SI for energy-momentum diagrams). The SI section also includes a comparison of spectra taken at different islands to validate the reproducibility of the results and the homogeneity of the sample.

The spectra of Figure 2a show a prominent feature at ~1.4 eV below  $E_{\rm E}$  that shifts down slightly to higher binding energies as the gold coverage increases. We identify this photoemission feature as having significant MoS2 valence band weight. Given a MoS<sub>2</sub> bandgap of 2.1-2.5 eV (roughly equivalent to the optical bandgap of 1.87 eV plus exciton binding energy), 45-47 the position of the valence band edge at  $\Gamma$  (which is slightly below that at K) bespeaks a largely undoped material. This observation affirms the high quality of the CVD MoS2 singlelayer film of this study, because the typical defects in the MoS, material, sulfur vacancies, act as n-type dopants. From the small value of the Fermi level shift  $\Delta E_{\rm F}$  away from the midgap position by only 0.1-0.3 eV and a straightforward application of the standard formula for the Fermi level shift  $\Delta E_{\rm F} = k_{\rm B} T \ln \frac{n_{\rm doping}}{n_{\rm intrinsic}}$  based on the Boltzmann constant ( $k_{\rm B}$ ), the temperature (T), and the defect-induced vs intrinsic charge carrier concentration  $(n_{\text{doping}}/n_{\text{intrinsic}})$ , we obtain a value of the defect-induced free carrier concentration far smaller than one per square micron. While this analysis omits carriers trapped by chemical passivation of the vacancies, it clearly points to a defect density in our samples that is so low that defects will not determine the overall material properties.

The small down-shift ( $\sim$ 200 meV) of the valence band edge with increased gold coverage suggests strong interactions between MoS<sub>2</sub> and the adlayer Au, and possibly very moderate additional n-type doping of the MoS<sub>2</sub> material as well. Analysis of the gold-induced Fermi-level shift ( $\Delta E_{\rm F}$ ) in a similar fashion as described above suggests that it reflects transfer of only a fractional charge per few-micron island. While a precise numeric analysis of the MoS<sub>2</sub> valence band peak position is precluded by the nontrivial background, it is clear that the majority of the shift in our experiments occurs even before a monolayer of gold is reached, suggesting the significance and chemical strength of the direct interaction between the gold adatoms and the substrate. This interaction—and its catalytic implications—will be the focus of the remainder of the manuscript.

Deposition of gold on single-layer MoS2 leads to a broad density of states (DOS) contribution at 3-4 eV below the Fermi level. Comparing the electronic structure of low coverages of gold on MoS2 to that of gold on graphene (Figure 2c), the absence of a well-defined and sharp gold 5d peak or doublet (dashed lines) is noticeable. Gold is known to aggregate into islands (Vollmer Weber Growth) on graphene and graphite, giving rise to extended gold aggregates of welldefined metallic band structure even at very low average coverage. 10,11 In contrast, both the rapid attenuation of the MoS<sub>2</sub> valence band peak with gold deposition in the data set of Figure 2a (a numerical analysis is shown in the SI) and the emergence of a broad gold 5d peak suggest the contrary for deposition onto MoS<sub>2</sub>. In further support of this interpretation, we performed scanning tunneling microscopy imaging of comparable gold coverages on cleaved (bulk) MoS<sub>2</sub> surfaces and find the formation of highly dispersed small islands. 48 Prior vibrational spectroscopy also suggests strong mechanical attachment in this material system.

### DISCUSSION

DFT modeling of the total density of states of gold aggregates of different sizes and shapes on a single layer thick sheet of MoS<sub>2</sub> provides insight into the strength of the Au-MoS<sub>2</sub> interaction at the root of the high dispersion and absence of aggregation. Figure 2b shows the Au 5d DOS of an individual gold atom as well as two flat and one bilayer gold clusters on MoS<sub>2</sub>. The calculations do not include the SiO<sub>2</sub>/Si substrate and, hence, do not describe any substrate-effects on the Fermi level position. A single gold atom exhibits a prominent Au 5dderived DOS contribution at a binding energy of ~2.5 eV (red arrow). As the size of the gold aggregate on MoS<sub>2</sub> increases, a number of additional 5d-derived DOS contributions emerge both at higher binding energy and at  $\sim$ 2.4 eV (blue arrow). The latter dwarves the former as the island size reaches 19 Au atoms (navy in Figure 2b). This trend can also be observed in the experimental data (Figure 2a), where a similar shift of peak intensity toward lower binding energy occurs as the Au coverage increases. Unfortunately, the multitude of substrate contributions prohibit numeric spectral deconvolution.

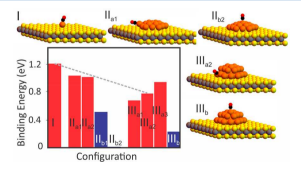
Our findings pose the question of the origin of aforementioned two spectral contributions—and how they emerge from Au-MoS<sub>2</sub> interactions. Figure 3 plots the spatial distribution of the dominant contribution to the local density of states (LDOS); volume elements, in which the LDOS at the higher binding energy dominates are colored in red, while volume elements in which the lower binding energy state dominates are colored in blue. The top panels show the LDOS distributions in the two energy intervals individually. In the case of the single atom, the higher binding energy state is in the surface plane whereas the, whereas the lower binding energy is of  $dz^2$  (a<sub>1</sub>) symmetry. The 19-atom cluster is held on the substrate by interactions at its edge leading to a very moderate curvature in the cluster similar to that found in the top layer of the bilayer cluster.

Islands of different sizes share in common that at their perimeter or at their bottom (in the case of the bilayer island), i.e., wherever the interaction with the MoS<sub>2</sub> is enhanced, the higher binding energy state dominates, and elsewhere the lower binding energy one. The cross-sectional views affirm that the higher binding energy state hybridizes significantly with the top sulfur layer of the substrate, whereas the lower binding energy state does not.

The gold d-band position and its hybridization  $^{21,50}$  have been determined to be a crucial descriptor of its catalytic activity. Hence, the ability of single-layer  $MoS_2$  to interact sufficiently strongly with gold to affect a major descriptor of its catalytic potential suggests that  $MoS_2$  may have a use wherever a corresponding shift of the Au 5d state is desired.

Computationally the resultant affinity of the Au clusters to adsorption of CO molecules depends very much upon Au cluster size and the site. The various CO adsorption sites investigated are indicated in Figure 3 by Roman numerals standing for the cluster size followed by labels a,b to indicate whether the location is one at which the higher or lower binding energy LDOS dominates, respectively.

The results (Figure 4) exhibit significant variation of the CO adsorption energy between different CO adsorption sites



**Figure 4.** Impact of substrate interactions: the bar graph shows the adsorption energy of a CO molecule at different locations on a single Au atom (I), on a single-layer Au cluster (II), and on a bilayer Au cluster (III). Site labels are defined in Figure 3 and examples of the adsorption geometries are shown. The color coding of the bars represents the dominance of the higher/lower (red/blue) binding energy LDOS as shown in Figure 3.

ranging from practically no chemical bonding ( $\sim$ 0.01 eV) in the center of the single-layer island ( $\rm II_{b2}$ ) to quite strong bonding (1.2 eV, about double that found on gold low-index surfaces) at the isolated gold atom and the perimeter sites of the clusters. The latter originates from a combination of

enhanced CO-gold and CO-substrate interactions in binding configurations strongly tilted toward the surface plane. In each case, we find that the locations on the cluster, in which the Au-MoS<sub>2</sub> substrate interactions rendered the higher binding energy Au 5d component dominant, the CO binding energy exceeded that of the other sites and also that of bulk gold surfaces. On average, we find that the enhancement of the gold-CO interaction diminishes with increasing cluster size (dashed lines in Figure 4) toward the typical binding energy of CO on Au of 0.6 eV (0.42 eV in our calculations) observed both for the bulk and for gas-phase clusters of increasing size. These results are also in agreement with the well-established inactivity of bulk gold and the findings of ref 21 for O<sub>2</sub> adsorbates.

The computational approach also allows us to compare the binding energy of CO molecules on gold clusters supported by MoS<sub>2</sub> to CO binding to unsupported gold clusters of the same shape. We find that in the configurations these clusters are held in by MoS<sub>2</sub>, they are also expected to exhibit strong binding in the gas phase. The SI provides a comparison.

#### CONCLUSIONS

Our combined experimental-computational investigation finds that single-layer MoS<sub>2</sub> islands can be prepared by scalable chemical vapor deposition techniques on a very thin oxide substrate where they offer an electronic structure closely resembling that of exfoliated MoS<sub>2</sub> and, most importantly, capable of interacting sufficiently with gold atoms so as to cause dispersion into small clusters. The resultant clusters are locked into a shape that is associated with undercoordination of edge gold atoms sustaining the substrate interaction and generating strong affinity for binding of small molecule species such as CO adsorbates at binding energies in excess of 1 eV, i.e., even at noncryogenic temperatures. Future work will explore whether such binding configurations can serve as initial states for catalytic activation of CO and other small-molecule species. The facile and conformal nature, in which MoS<sub>2</sub> can be deposited by CVD onto a broad range of oxide substrate suggests great versatility of these findings.

# ASSOCIATED CONTENT

### S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b07632.

Energy-momentum diagrams showing the dispersion of the single-layer  $MoS_2$  film; spectroscopic data validating lateral homogeneity of the film; spectroscopic data validating the gold deposition thickness; and quantitative comparison of computational CO adsorption energies (PDF)

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Notes

The authors declare no competing financial interest.

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